

Photoconversion Mechanism of a Green/Red Photosensory Cyanobacteriochrome AnPixJ: Time-Resolved Optical Spectroscopy and FTIR Analysis of the AnPixJ-GAF2 Domain

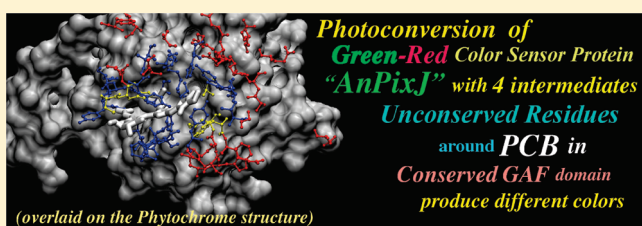
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ABSTRACT: The photoconversion mechanism of a green/red sensory cyanobacteriochrome AnPixJ was studied. The phycocyanobilin-binding second GAF domain of AnPixJ of *Anabaena* sp. PCC 7120 was expressed in *Escherichia coli* cells. The His-tagged AnPixJ-GAF2 domain exhibited photoconversion between the green- and red-absorbing forms, APg₅₄₃ and APr₆₄₈, respectively. We detected four intermediate states in the photocycle between them, as follows: APr₆₄₈ → red light → APr₆₄₈* → (with a rise time constant τ_r of <100 ns) R1_{650–80} (with a decay time constant τ_d of <1 μ s) → R2₆₁₀ (τ_d = 920 μ s) → APg₅₄₃ → green light → APg₅₄₃* → (τ_r < 50 ns) G1₅₇₀ (τ_d = 190 μ s) → G2₆₃₀ (τ_d = 1.01 ms) → APr₆₄₈. These intermediates were named for their absorption peak wavelengths, which were estimated on the basis of the time-resolved difference spectra and global analysis of the time courses. The absorption spectrum of APr₆₄₈ resembles that of the Pr form of the phytochrome, while all the other states showed peaks at 530–650 nm and had wider bandwidths with smaller peak amplitudes. The fastest decay phases of fluorescence from APr₆₄₈* and APg₅₄₃* gave lifetimes of 200 and 42 ps, respectively, suggesting fast primary reactions. The APg₅₄₃-minus-APr₆₄₈ difference FTIR spectrum in an H₂O medium was significantly different from those reported for the Pfr/Pr difference spectra in phytochromes. Most of the peaks in the difference spectrum were shifted in the D₂O medium, suggesting the high accessibility to the aqueous phase. The interactions of the phycocyanobilin chromophore with the surrounding amino acid residues, which are fairly different from those in the GAF domain of phytochromes, realize the unique green/red photocycle of AnPixJ.



Color perception is vital for photosynthetic organisms. Phytochromes and bacteriophytochromes have been known to regulate light acclimation of plants and bacteria as red/far-red photoreceptors.^{1–7} They contain GAF (cGMP-phosphodiesterase/adenylate cyclase/formate hydrogen lyase transcription activator FhlA) domains and bind bilin pigments as chromophores. Their red- and far-red-absorbing forms (Pr/Pfr forms) are well-known to be converted to each other upon red/far-red light absorption.^{1,2} Multiple types of flavoproteins, such as cryptochrome, LOV, and BLUF domain proteins and photoactive yellow protein, are known to function as the blue light sensor.^{8–14} Only the retinal families have been known as the green light sensor until recently, in a cyanobacterium *Anabaena* (*Nostoc*) sp. PCC 7120.¹⁵ However, genomic analyses of cyanobacteria recently indicated another type of light sensor, cyanobacteriochromes.^{16–25} Cyanobacteriochromes have GAF domains, bind bilin chromophores, and sense green/red or blue/green light.^{16–25} The response to the green light, thus, seems to be a common feature of cyanobacteria.

In *Synechocystis* sp. PCC 6803 (*Synechocystis* 6803), a blue/green cyanobacteriochrome SyPixJ (PixJ1) protein^{26,27} regulates the phototaxis of cells together with a BLUF photoreceptor, SyPixD.⁹ Photoconversion of another blue/green sensory cyanobacteriochrome,

TePixJ, of *Thermosynechococcus elongatus* BP-1 has also been reported.^{18,19,25} As the green/red sensors, cyanobacteriochrome CcaS proteins of *Synechocystis* 6803 and *Nostoc punctiforme* are reported to regulate the chromatic acclimation of the photosynthetic light-harvesting system.^{23,28} A cyanobacteriochrome, AnPixJ, of *Anabaena* sp. PCC 7120 also senses green/red light and is deduced to regulate the phototaxis of cells.²¹

The AnPixJ protein is composed of four GAF domains and one MCP domain. The second GAF domain (designated AnPixJ_{GAF2} hereafter) binds phycocyanobilin [PCB (Figure 1C)] when expressed in *Escherichia coli* together with a pigment-synthesizing system.²¹ We used the His-tagged AnPixJ_{GAF2} domain expressed in *E. coli* in this study. The purified AnPixJ_{GAF2} shows the green- and red-absorbing forms with peaks at 543 and 648 nm (designated APg₅₄₃ and APr₆₄₈, respectively, hereafter), as shown in Figure 1A. APr₆₄₈ gives a spectrum of the red absorption band, which resembles the absorption spectrum of the Pr form of the phytochrome that gives a peak at 667 nm with an ~50 nm bandwidth. APr₆₄₈ and APg₅₄₃ interchange with each

Received: November 10, 2010

Revised: June 15, 2011

Published: June 29, 2011

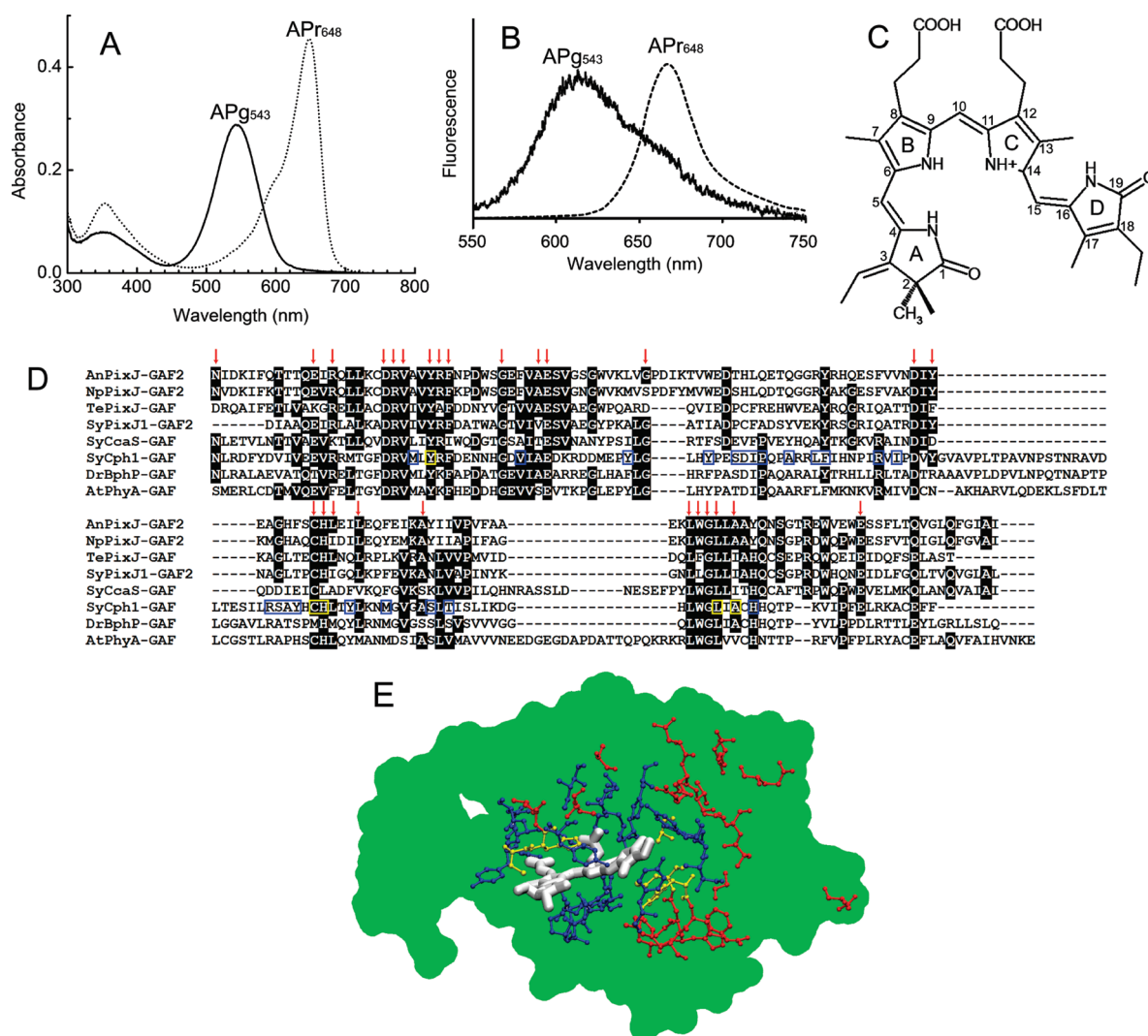


Figure 1. (A) UV–visible absorption spectra of APg₅₄₃ (—) and APr₆₄₈ (···) forms of AnPixJ_{GAF2}. (B) Fluorescence spectra of APg₅₄₃ (—) and APr₆₄₈ (---) forms excited at 360 and 500 nm, respectively. (C) Molecular (ZZZssa) structure of PCB. (D) Alignment of amino acid residues created with eight GAF domains of PixJs, phytochromes, and bacteriophytochromes. Residues identical among more than four GAF domains are denoted with a black background. Blue and yellow boxes represent residues forming chromophore-binding pockets in SyCph1. The residues inside blue and yellow boxes are unconserved and conserved, respectively, between AnPixJ_{GAF2} and SyCph1. The residues within 5 Å of the chromophore were picked up from the structures of Pr forms of SyCph1 (Protein Data Bank entry 2VEA). Red arrows indicate the residues conserved between AnPixJ_{GAF2} and SyCph1. (E) Spatial arrangements of PCB and amino acid residues on the GAF domain of SyCph1 in Pr form (Protein Data Bank entry 2VEA) that are conserved and unconserved between SyCph1-GAF and AnPixJ_{GAF2}. The moiety drawn as a white stick model represents the PCB chromophore, and the ball-and-stick model represents the amino acid residues shown in the homology analysis in panel D. The green part represents the whole GAF domain of SyCph1 shown in panel D. Yellow and red portions are conserved residues in contact and not in contact with PCB, respectively. Blue portions are unconserved residues in contact with PCB. Names of proteins and sequences are as follows: An, *Anabaena* (*Nostoc*) sp. PCC 7120; Np, *N. punctiforme* ATCC 29133; Te, *T. elongatus*; Sy, *Synechocystis* sp. PCC 6803; Dr, *D. radiodurans*; At, *Arabidopsis thaliana*; AnPixJ, All1069 (NP_485112); NpPixJ, NpR6012 (ZP_00110373); TePixJ, Tll0569 (NP_681359); SyPixJ1, Sll0041 (NP_442716); SyCcaS, Sll1473–1475 (ABI83649); SyCph1, Slr0473 (Q55168); DrBphP, DR_A0050 (Q9RZA4); AtPhyA, AT1G09570 (NP_172428).

other by the absorption of red and green light, respectively. The APg₅₄₃ form is stable for several hours in the dark and converts to the APr₆₄₈ form after a very long dark adaptation period, indicating that APr₆₄₈ is a resting form.²¹ Narikawa et al.²¹ proposed a change of pigment color to reflect the isomerization of the D-ring of PCB based on the acidic urea denaturing experiment.

The amino acid sequence of AnPixJ_{GAF2} shows homology to those of the GAF domains of canonical phytochromes or bacteriophytochromes, as shown in Figure 1D. The residues in the chromophore-binding pockets, which are in blue and yellow

boxes in Figure 1D, are highly conserved between the sequences of cyanobacterial phytochrome Cph1 (*Synechocystis* 6803) and bacteriophytochrome DrBphP (*Deinococcus radiodurans*). Among the 27 residues that are in contact with (within 5 Å of) the PCB chromophore in the Cph1 structure, 17 residues are conserved in DrBphP. On the other hand, only five residues are common between the sequences of AnPixJ_{GAF2} and Cph1 (yellow boxes, Figure 1D).

This suggests different environments and binding sites of chromophores in these proteins (see Figure 1E, too). In this

study, we extended the analysis of the photoconversion between APr_{648} and APg_{543} , which was reported briefly in our previous report,²¹ and identified the reaction times and spectral shapes of four intermediate states. We compared them to those of phytochromes to determine the mechanism of the color change and sensing of $\text{AnPixJ}_{\text{GAF2}}$. It is interesting that almost the same bilin pigments in GAF domains in the two groups of proteins undergo very different color changes.

In the $\text{Pr} \rightarrow \text{Pfr}$ photoconversion of phytochromes, three major intermediates have been proposed.² A sequential transition from the excited- Pr to lumi-R (with an absorption peak at 688 nm), meta-Ra (663 nm), and meta-Rc (725 nm) has been proposed in the $\text{Pr} \rightarrow \text{Pfr}$ conversion.² The time-resolved spectroscopy of the native oat phytochrome has revealed five kinetic components in the $\text{Pr} \rightarrow \text{Pfr}$ conversion, which are named lumi-R1 , lumi-R2 , meta-Ra1 , meta-Ra2 , and meta-Rc .²⁹ In the $\text{Pfr} \rightarrow \text{Pr}$ conversion, lumi-F (670 nm, 320 ns), meta-Fa (656 nm, 265 μs), and meta-Fb (660 nm, 5.5 ms) are proposed as intermediates in a sequential reaction scheme.³⁰

Vibrational spectroscopy has also been used to study phytochromes in combination with isotope labeling and density functional theory calculations. The Pfr/Pr -type phytochromes showed characteristic difference FTIR spectra.³¹ The phytochrome of the non- Pfr/Pr type, such as a bacteriophytochrome RbBphP3 that binds biliverdin (BV) and converts between the Pr (705 nm) and Pnr (645 nm) forms, exhibited somewhat different IR spectra.⁷ Ultrafast mid-infrared spectroscopy on cyanobacterial Cph1 revealed structural changes in the picosecond time range and proposed the $15Z \rightarrow E$ isomerization of the chromophore for the $\text{Pr} \rightarrow \text{lumi-R}$ conversion.³² Time-resolved spectroscopy, Raman spectroscopy, and quantum chemical calculations on phytochromes also predicted the changes in the protonation state during the Pr to Pfr photoconversion.^{33–35}

In this study, we identified the multiple intermediate states in the green/red photoconversion process of $\text{AnPixJ}_{\text{GAF2}}$. The intermediate states in the photoconversion of cyanobacteriochromes have not been reported yet, except for our preliminary results for $\text{AnPixJ}_{\text{GAF2}}$.²¹ The results of this study identified multiple intermediates in the photocycle of $\text{AnPixJ}_{\text{GAF2}}$ and suggested the structural changes of the protein and chromophore that are fairly different from those in the photocycle of phytochromes.

MATERIALS AND METHODS

Protein Expression and Purification. His-tagged $\text{AnPixJ}_{\text{GAF2}}$ was purified as described previously.²¹ *E. coli* cells that contain the expression vectors of His-tagged $\text{AnPixJ}_{\text{GAF2}}$ and chromophore expression^{21,36} were harvested by centrifugation and frozen at -80°C . The collected cells were resuspended in a 20 mM HEPES-NaOH buffer (pH 7.5) and disrupted with a French pressure cell. The disrupted cell debris was removed by centrifugation, and after imidazole had been added to the supernatant, the solution was loaded onto a Ni affinity column (HiTrap Chelating HP, GE Healthcare, Piscataway, NJ). The column was washed with a washing medium containing 20 mM HEPES-NaOH (pH 7.5), 100 mM NaCl, and 100 mM imidazole and then eluted using a step gradient of imidazole in an elution medium. The obtained protein solution was dialyzed against a reaction medium containing 20 mM HEPES-NaOH (pH 7.5), 100 mM NaCl, and 10% glycerol and stored frozen until use.

Identification of Photoconversion Intermediates by Laser Spectroscopy. The transient time course of the absorption

change induced by flash excitation was measured with a home-built split beam spectrophotometer as described previously³⁷ with an instrument response time of 30 μs . The sample solution in an acrylic cuvette (optical path length of 10 mm) was excited by a 10 ns laser flash from SHG of a Nd:YAG laser (at 532 nm, 10 mJ; Quanta-Ray LAB-130-10TH, Spectra Physics, Santa Clara, CA) for the measurement of the $\text{APg}_{543} \rightarrow \text{APr}_{648}$ conversion or Xe flash light (10 μs duration; Sugawara Laboratories, Inc., Kawasaki, Japan) for the measurement of the $\text{APr}_{648} \rightarrow \text{APg}_{543}$ conversion at wavelengths of >620 nm through a 620 nm long-pass filter. To ensure the photoreaction from the fully converted APg_{543} or APr_{648} state, the photoproduct after each flash excitation was reversed by illumination with a 530 or 627 nm LED (LXHL-LM5C or LXHL-LD3C, respectively; Philips Lumileds Lighting, San Jose, CA). The intensity and duration of the illuminations were 16 mW/cm^2 and 1 s for the 530 nm LED and 14 mW/cm^2 and 0.25 s for the 627 nm LED, respectively. Curve fittings and global analysis of time courses were performed using Microcal Origin 6.0 (Microcal Software, Inc., Northampton, MA).

The photoconversions were also studied from shorter time ranges by measuring the time-resolved spectra with a multi-channel spectrophotometer equipped with an image intensifier-gated diode array (Princeton Instruments, Trenton, NJ) or an image intensifier-gated CCD (Instaspec ICCD detector model 77193-5, ORIEL Instruments, Stratford, CT) with a time resolution of 20 or 50 ns, respectively, as reported previously.^{10,37} The 10 ns excitation flash at 620 nm (at ~ 1 mJ at a repetition rate of 0.03 Hz) was obtained from a solid-state dye laser (CF171 Astra, Solar TII, Minsk, Belarus), which was pumped by a 532 nm laser flash from a Nd:YAG laser (LS-2135, Solar TII). The excitation at 532 nm was conducted with the Nd:YAG laser. Time-resolved difference absorption spectra were monitored by a probing light from a 10 μs Xe flash light. The probing light that passed through the sample was diffracted in a spectrograph (MS157, ORIEL Instruments) and detected by a gated diode array or a gated ICCD detector that was controlled by a timing generator (DG535, Stanford Research Systems, Sunnyvale, CA). The diode array or the ICCD was activated for 20 or 50 ns in concert with the probing flash at varied times with respect to the laser peak time. Photoproducts formed by each excitation laser flash were, then, reverted by illuminating LED light for a few seconds following each excitation—measurement cycle as described for the time course measurement.

Measurement of Fluorescence Spectra and Lifetimes. Steady-state fluorescence spectra of APg_{543} and APr_{648} forms were measured with a fluorescence spectrometer (FL4500, Hitachi High-tech, Tokyo, Japan). Measurement of fluorescence lifetimes was conducted with a streak camera system (Hamamatsu 4334, Hamamatsu Photonics, Inc., Hamamatsu, Japan) as described previously.^{38,39} APg_{543} or APr_{648} forms of an $\text{AnPixJ}_{\text{GAF2}}$ solution were excited with a SHG of a Ti:Sapphire laser at 355 nm (at 50 μW , 150 fs pulse width; Mai Tai, Spectra Physics) or a 630 nm diode laser (at 0.65 μW , 50 ps pulse width; Hamamatsu Photonics, Inc.) fired at 40 or 1 MHz, respectively, and the photons emitted from the samples were detected in a photon counting mode. The excitation beam intensities at $\sim 10^{-18}$ mol/pulse are weak enough to excite only a small fraction of the sample at a low concentration of ~ 5 nmol/ μL . Therefore, fluorescence is expected to be emitted from either APg_{543} or APr_{648} , and not from their doubly excited (excited intermediate) forms. We did not notice significant contributions of the intermediate fluorescence in the time-resolved spectra of

the slow decay components. The instrument response times were 21 and 50 ps for the 355 and 630 nm excitations, respectively. Acquisition of data was stopped every 5 min, and the samples were exposed to illumination with a 660 nm laser diode (1 mW/cm²; HL6535MG, Opnext, Inc., Fremont, CA) or the 530 nm LED (0.7 mW/cm²; LXHL-LMSC) for 1 min to fully convert the photoproducts into the initial states, in the whole measuring time of 0.5–1 h. The obtained fluorescence decay wavelength two-dimensional data were analyzed with a global analysis program as described previously.^{38,39}

FTIR Measurement. The stock solution of AnPixJ_{GAF2} (at ~1 mL at ~8 μ M) was concentrated by centrifugal filtration (Vivaspin 500, 10000 molecular weight cutoff, Sartorius). The sample was diluted 10-fold with either H₂O or D₂O medium containing 20 mM HEPES and 100 mM KCl at pH or pD 7.5 and again concentrated by centrifugal filtrations. The dilution and concentration cycle was repeated two or three times to obtain the buffer-exchanged solution of AnPixJ_{GAF2} at the final concentration of 0.2 mM. Samples were held at 0–10 °C during the preparation. A 5 μ L aliquot of the AnPixJ_{GAF2} solution was placed between two CaF₂ disks, and the disks were sealed with silicon grease and then illuminated with either a 520 or 645 nm LED to fully convert to the APr₆₄₈ or the APg₅₄₃ form, respectively. Light-induced difference FTIR spectra were recorded with a Bruker IFS66/S spectrometer equipped with a liquid nitrogen-cooled MCT-A detector as described previously,³¹ which gives the spectral resolution of 4 cm⁻¹ with an accuracy of ± 1 cm⁻¹. After a background spectrum (an average of 400 interferograms measured for 55 s) during the 520 nm illumination had been recorded, the illumination light was switched to a 645 nm frequency, and after a 5 s delay, the sample spectrum (400 interferograms) was recorded to produce an APg₅₄₃-minus-APr₆₄₈ difference spectrum. A new background was then taken, and the light was switched back to the 520 nm LED once again. After a 5 s delay, a sample spectrum was taken during the 520 nm illumination to produce an APr₆₄₈-minus-APg₅₄₃ difference spectrum. To improve the signal-to-noise (S/N) ratio, the illumination cycle was repeated 250 times, and the obtained APg₅₄₃-minus-APr₆₄₈ spectra and the inverse of the APr₆₄₈-minus-APg₅₄₃ spectra were averaged to produce a final APg₅₄₃-minus-APr₆₄₈ spectrum. The illumination cycle was automated and synchronized with IR scanning by using a laboratory-made electric circuit. The sample was fully active during the IR measurement at 15 °C.

RESULTS

Photoconversion between APg₅₄₃ and APr₆₄₈ Forms Induced by Continuous Illumination. As shown in Figure 1A, AnPixJ_{GAF2} is interconverted between the green-absorbing APg₅₄₃ and the red-absorbing APr₆₄₈ forms upon the absorption of green and red light, respectively. The conversion under continuous illumination gave isosbestic points at 582 and 452 nm, indicating the rapid interchange between these two states. The absorption band of APr₆₄₈ has a higher peak amplitude with a narrower bandwidth giving a clear shoulder. Therefore, the conversion includes a process to shift the peak and to change the spectral shape, suggesting a large change in the molecular structure of PCB inside the protein. The spectral shape of APr₆₄₈ resembles that of the Pr form of the phytochrome. The fluorescence spectra of APg₅₄₃ and APr₆₄₈ forms exhibited emission

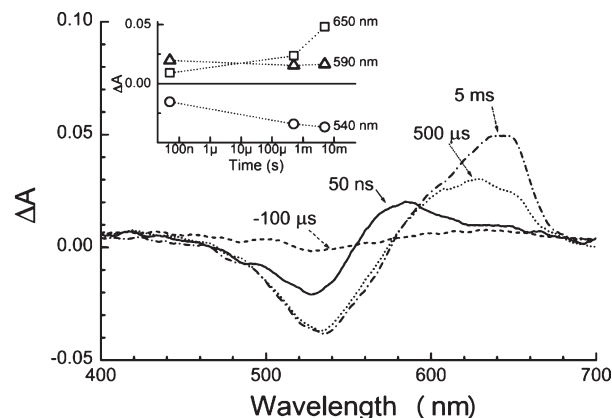


Figure 2. Time-resolved difference absorption spectra measured during the photoconversion from APg₅₄₃ to APr₆₄₈ states of AnPixJ_{GAF2}. Spectra were recorded with a gated diode array system at delay times of -100μ s (before the laser excitation), 50 ns, 500 μ s, and 5 ms with respect to the time of excitation with a 10 ns, 532 nm laser flash. The inset shows the amplitude of the absorption change at each wavelength plotted vs the logarithm of time.

peaks at 610 and 670 nm by excitation at 360 and 500 nm, respectively (Figure 1B).

Detection of Intermediates in the Photoconversion from APg₅₄₃ to APr₆₄₈. Time-resolved difference absorption spectra were recorded with a gated diode array spectrograph to identify the reaction intermediates in the photoconversion reaction from APg₅₄₃ to APr₆₄₈ after excitation with a 10 ns laser flash at 532 nm (Figure 2). To avoid the accumulation of photoproducts during the repetitive flash excitations, the sample was illuminated with a red LED light between the excitation–measurement periods. The difference spectrum at 50 ns showed a negative peak at 530–540 nm and a positive peak at 590 nm. The difference spectrum at 500 μ s showed a larger negative peak at 530–540 nm with a higher positive peak at 630 nm. The difference spectrum at 5 ms showed a similar extent of a negative peak at 540 nm and a larger positive peak at 640 nm. This difference spectrum is similar to the APr₆₄₈-minus-APg₅₄₃ difference spectrum induced by continuous illumination, indicating the formation of APr₆₄₈ from APg₅₄₃ at 5 ms. The inset shows the time-dependent changes in the amplitudes at each wavelength. These time courses and difference spectra, which differ from each other, suggested at least two intermediate states in the conversion from APg₅₄₃ to APr₆₄₈.

Time courses of the absorption changes at each wavelength were also monitored continuously after excitation with the 532 nm laser flash. In Figure 3A, the extents of absorption changes at 565, 610, and 650 nm were plotted versus the logarithm of time after laser excitation over the wide time range of 0.1–100 ms. Time courses with low noise and high time resolution were obtained by the use of 10 ns, 532 nm laser excitation, which allowed the selective excitation of APg₅₄₃. Each time course suggested multiple intermediate forms. Similar time courses measured every 10 nm from 450 to 750 nm (not shown) were, then, fitted as the sum of three exponentially decaying components and one nondecaying component. The absorption spectrum and reaction time constant of each intermediate were then assumed by a global analysis of all the time courses. The fastest kinetic component was resolved to appear with a time constant of 20 μ s (i.e., within the response time of the apparatus). This component, which partially includes the contribution from

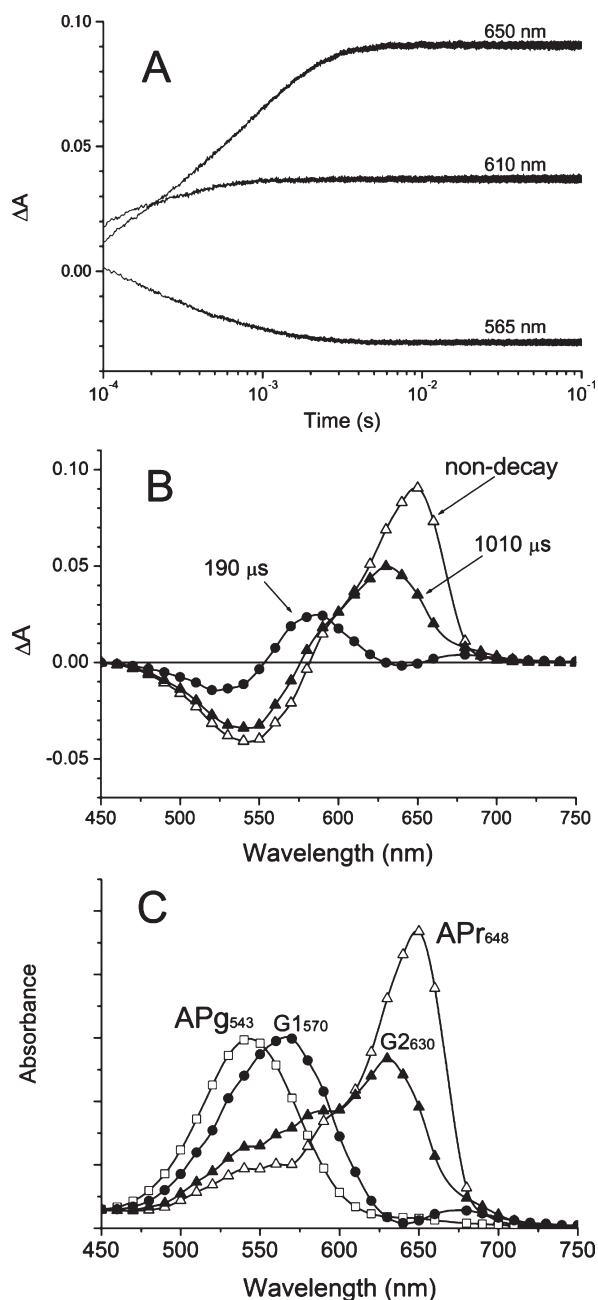
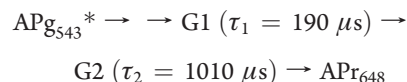


Figure 3. (A) Time courses of the absorption change at 565, 610, and 650 nm induced by the laser excitation of APg₅₄₃. (B) EADS of each intermediate state calculated from the set of time courses measured each 10 nm over the range of 450–750 nm. Each EADS was calculated by assuming the sequential formation of each intermediate in a single-exponential decay process. Filled circles and triangles represent the EADS with 190 μ s (G1) and 1010 μ s (G2) decay time constants, respectively. Empty triangles represent the photoproduct nondecaying component. (C) The absorption spectrum of each intermediate state was estimated by adding each EADS sequentially to the absorption spectrum (\square) of APg₅₄₃ with appropriate normalizations as described in the text. Filled circles, filled triangles, and empty triangles represent the estimated absorption spectra of the 190 μ s (G1), 1010 μ s (G2), and nondecay (APr₆₄₈) components, respectively.

the fluorescence from the excited state of APg₅₄₃, was not further analyzed. The two slower components, which had decay time constants of 190 and 1010 μ s, were further analyzed.

We analyzed the amplitudes of the 190 μ s (τ_1) and 1010 μ s (τ_2) decay phases at all wavelengths to obtain the absorption spectra of reaction intermediates. Figure 3B shows the evolution-associated difference spectra (EADS) obtained by assuming a simple scheme and rate equations, in which sequential reactions take place as follows



where τ_1 and τ_2 represent the lifetimes of two intermediate states. The EADS with a τ_1 of 190 μ s had a negative peak at 520 nm and a positive one at 590 nm (the intermediate state is designated G1). The spectrum is almost the same as the time-resolved spectrum at 50 ns in Figure 2 and indicates the formation of the G1 state from APg₅₄₃ within 50 ns of laser excitation and its conversion to the second intermediate (G2) with a time constant (τ_1) of 190 μ s (see Discussion).

The second EADS with a τ_2 of 1010 μ s showed a negative peak at 540 nm and a positive peak at 630 nm with a shoulder at \sim 590 nm. The difference spectrum indicates the red shift of the peak wavelength of the absorption band from 543 nm in APg₅₄₃ to 590 nm in G1 and then to 630 nm in G2. The difference spectrum of the nondecaying component showed a negative peak at 540 nm and a positive peak at 650 nm. It was almost the same as that of the APr₆₄₈-minus-APg₅₄₃ difference spectrum obtained at the steady state. These results indicate that the transition from G2 to APr₆₄₈ takes place with a τ_2 of 1010 μ s.

The absorption spectra of G1 and G2 were estimated by adding each EADS spectrum to the (steady-state) absorption spectrum of APg₅₄₃ by using an adequate normalization factor that was calculated from the comparison of “the nondecaying component of EADS” and “the APr₆₄₈-minus-APg₅₄₃ difference spectrum” (Figure 3C). The absorption spectra of APg₅₄₃, G1, G2, and APr₆₄₈ showed peaks at 543, 570, 630, and 648 nm, respectively. It is clear that the photoconversion proceeds with the hopping of peak wavelengths from 543 nm to the red until 648 nm is reached in \sim 0.1 s, together with the changes in the conformation of the protein. The spectra of APg₅₄₃ and G1 show similar bandwidths and peak amplitudes with 27 nm-shifted peak wavelengths. The estimated G2 spectrum gave a peak at 630 nm with a similar or slightly smaller peak amplitude and a wider bandwidth. The spectral feature of G2 might also contain small contributions from G1 and APr₆₄₈ because the wide bandwidth of G2 extended to the spectral ranges of these states. This may happen if the actual decays do not obey single exponentials. On the other hand, it is clear that APr₆₄₈ has a higher peak amplitude and a narrower bandwidth compared to the other states. This suggests a significant change in the chromophore structure in the conversion from G2 to APr₆₄₈.

Detection of Intermediates in the Photoconversion from APr₆₄₈ to APg₅₄₃. Time-resolved difference absorption spectra were also recorded in the photoconversion from APr₆₄₈ (Figure 4). We measured the difference spectrum at each indicated time after the 20 ns, 620 nm laser-flash excitation by using the gated ICCD to eliminate the effect of flash-induced strong fluorescence from APr₆₄₈. After each excitation–measurement period, photoproduct APg₅₄₃ was converted back to APr₆₄₈ via illumination of the green 530 nm LED light for 1 s to suppress the accumulation of photoproducts.

One hundred nanoseconds after the laser excitation of APr₆₄₈, we obtained a difference spectrum with a positive peak at 680 nm (indicated by the arrow marked a in Figure 4) and a large negative peak at 650 nm as reported previously.²¹ The negative peak

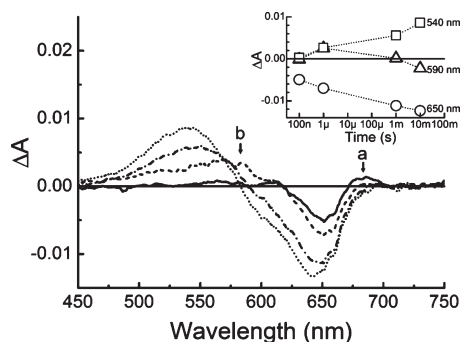


Figure 4. Time-resolved difference absorption spectra measured at 100 ns (—), 1 μ s (---), 1 ms (····), and 10 ms (· · · ·) after laser excitation at 620 nm on the APr₆₄₈ form of AnPixJ_{GAF2}. Arrows marked a and b denote the positions of apparent peaks at 680 and 570 nm, respectively. The inset shows the amplitude of the absorption change at each wavelength plotted vs the logarithm of time.

represents the bleaching of the ground state of the APr₆₄₈ band. The positive peak suggests the formation of a new intermediate, R1, which represents the formation of either a red-shifted band with a peak at 682 nm, as indicated by arrow a, or an intermediate at almost the same peak wavelength with a lower amplitude and wider bandwidth extended to the red side as discussed further below. The difference spectrum recorded at 1 μ s had no 680 nm positive peak and showed a broad positive peak at ~570 nm (indicated by arrow b), suggesting the formation of the second intermediate (R2) from R1 (Figure 4). At a delay time of 1 ms, the amplitude of the negative band around 650 nm increased more, and the positive peak was further blue-shifted to 550 nm. Ten milliseconds after excitation, the positive band showed a peak at 540 nm, indicating the conversion to the final APg₅₄₃ form. The inset time courses in Figure 4 indicate that an early intermediate R1 with a peak or shoulder at 680 nm is formed from APr₆₄₈ within 100 ns of the laser excitation and the second intermediate R2 is then formed within 1 μ s.

We also measured the time course of the absorption change of APr₆₄₈ induced by the excitation with a 10 μ s xenon flash (>620 nm) in the 100 μ s to 100 ms time range at each wavelength (Figure 5A). The large noise levels mainly came from the use of a fast response amplifier and the attenuated intensity of a probing light with the use of a monochromator to eliminate the broad and long scattering light and fluorescence induced by the xenon flash, the intensity of which was also low because of the limitation of the experimental setup for this measurement. The averaging and smoothing of abundant data points by an appropriate program, however, allowed detailed analysis of the time courses, especially over the longer time range. The whole process of photoconversion was completed within 5 ms of flash excitation. At 510 nm, the excitation induced only a small increase at the beginning and then gradually increased. At 580 nm, the instantaneous increase was larger, and the extent then gradually decreased.

We measured similar time courses each 10 nm over a range of 450–750 nm and calculated the EADS after analyzing each time course as a sum of decaying components that appear in a series. The analysis produced the difference absorption spectrum of an intermediate component (R2), which decays with a time constant of 920 μ s, together with the nondecaying one, which corresponds to the formation of APg₅₄₃, as shown in Figure 5B. The analysis of the time courses also indicated the formation of an R1 intermediate before R2, although the reaction time from

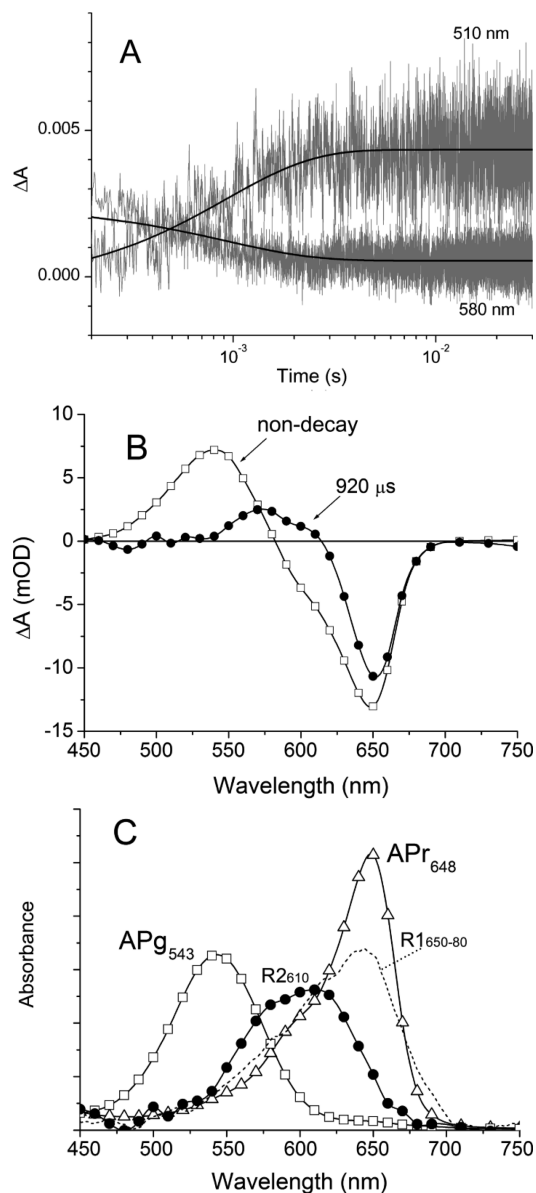
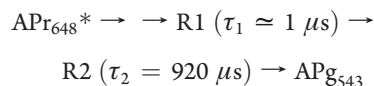


Figure 5. (A) Time courses of absorption change at 580 and 610 nm after the excitation of APr₆₄₈ with the red xenon flash. (B) EADS calculated from the time courses every 10 nm, like those in panel A, at 450–750 nm by assuming the sequential single-pathway scheme. Filled circles and empty squares represent the 920 μ s decay component (R2) and the nondecaying component, respectively. (C) Absorption spectra of intermediates estimated by adding each difference spectrum or EADS to the steady-state absorption spectrum of APr₆₄₈ (Δ). The dotted line represents the tentative absorption spectrum of R1. Filled circles and empty squares represent the estimated absorption spectra of the 920 μ s decay (R2) and nondecaying (APg₅₄₃) components, respectively.

R1 to R2 within 10 μ s could not be determined precisely because of the disturbance from the fluorescence excited by the long tail of the xenon flash. We used the time-resolved spectra in Figure 4 as well for the construction of the whole reaction process. On the basis of the results in Figures 4 and 5A, we assumed that the whole reaction process proceeds sequentially as follows.



The absorption spectra of intermediates R1 and R2 (dashed line and filled circles, respectively, in Figure 5C) were calculated by adding the difference spectrum/EADS to the steady-state APr₆₄₈ spectrum after adequate normalization. The spectrum of R1, which was estimated on the basis of the time-resolved spectrum in Figure 4, represents the results in Figure 5 well. The simulation result gave a spectrum with a peak amplitude lower than that of APr₆₄₈ with the shoulder extended to 680 nm rather than a spectrum with a red-shifted peak at 680 nm, although the result is still tentative. The spectrum of R2 is significantly blue-shifted to 610 nm. The peak amplitude of the R2 band after normalization is significantly lower than that of APr₆₄₈ and is close to that of APg₅₄₃. These results suggest that R1 has a lower peak amplitude at ~648 nm with a bandwidth wider than that of APr₆₄₈. The peak wavelength then shifted to 610 nm in R2 and to 543 nm in APg₅₄₃, showing similar peak amplitudes. The simulated absorption spectra of R1 and R2 intermediates are different from those of G1 and G2, suggesting different changes in chromophore configuration in the forward and backward photoconversions between APr₆₄₈ and APg₅₄₃.

Fluorescence Lifetimes of APr₆₄₈ and APg₅₄₃. The fluorescence spectra of APg₅₄₃ and APr₆₄₈ forms in Figure 1B showed emission peaks at 610 and 670 nm, respectively. To estimate the rates of the initial photoconversions from the excited states of APr₆₄₈ (APr₆₄₈^{*}) and APg₅₄₃ (APg₅₄₃^{*}), we measured the fluorescence lifetimes by using a streak camera system. We measured the wavelength–lifetime two-dimensional image of fluorescence after the excitation of APr₆₄₈ at 630 nm with a 50 ps diode laser pulse or APg₅₄₃ at 355 nm with a 0.2 ps pulse from a Ti:sapphire laser. A global analysis of the data points in different wavelength–time regions could resolve the decay time courses of multiple emission components even under the mixing conditions, as reported previously.^{38,39}

The fluorescence from APr₆₄₈ gave decay times of 200 ps (70% in amplitude) and 1000 ps (30%). APg₅₄₃, on the other hand, showed faster decay times of 42 ps (60%) and 610 ps (40%), as listed in Table 1. The time resolutions of the measurement of the APr₆₄₈ and APg₅₄₃ fluorescence were 50 and 21 ps, respectively. We assume the decay time constants of the major fast components in both states to be related to the photoconversion reactions and those of the slower components to be derived either from a heterogeneity or back reaction from the primary intermediates. The fluorescence of the Pr form of the phytochrome is reported to give two or more kinetic components,^{40,41} in which the fastest 30–60 ps component depended on the viscosity of the medium.⁴⁰ Biphasic decay was interpreted either by the heterogeneity of the sample or the fluorescence (delayed fluorescence) from the light-formed intermediate because of the reversibility of the reaction.⁴⁰

Recent studies by the ultrafast transient absorption measurement of the bacteriophytochrome RbBphP3 estimated the fast quenching process from Pr^{*} to occur with a 350 ps time constant because of the proton transfer–back-shuttle reaction between the chromophore (BV) and the amino acid, based on the observation of the slower rate in the D₂O medium.⁴² The fast fluorescence lifetimes of APr₆₄₈^{*} and APg₅₄₃^{*} may represent similar quenching mechanisms. The question of whether APr₆₄₈^{*} and APg₅₄₃^{*} are converted to R1 and G1 directly with time constants of 200 and 42 ps, respectively, or via some other early intermediates that cannot be detected at 50–100 ns by the present measurement of the absorption change remains to be answered.

Table 1. Fluorescence Lifetimes of APr₆₄₈^{*} and APg₅₄₃^{*}^a

fluorescence-emitting form (fluorescence peak wavelength)	lifetime (initial amplitude, relative)		
APr ₆₄₈ [*] (670 nm)	200 ps (0.7)	1000 ps (0.3)	
APg ₅₄₃ [*] (610 nm)	42 ps (0.6)	610 ps (0.4)	
Pr [*] (680 nm) ^b	39.6 ps (0.90)	167 ps (0.09)	1145.0 ps (0.01)

^a The lifetimes and relative initial amplitudes of decay components were calculated by the global analyses of wavelength–time two-dimensional images of the fluorescence as described in Materials and Methods.

^b Pr of 124 kDa oat phytochrome measured in potassium phosphate buffer (pH 7.8).⁴⁰

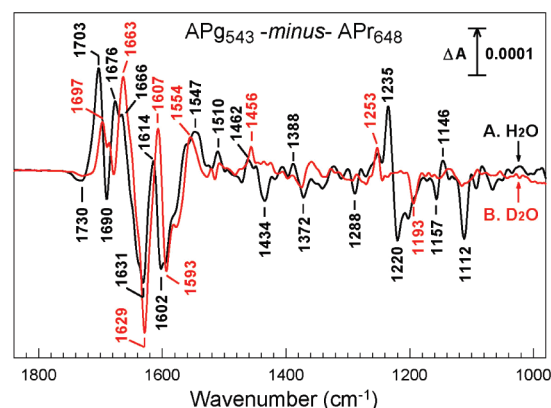
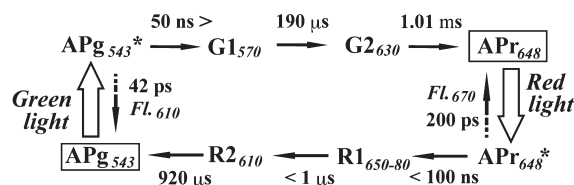


Figure 6. APg₅₄₃-minus-APr₆₄₈ difference FTIR spectra of AnPixJ_{GAF2}. Light-induced difference spectra of AnPixJ_{GAF2} were recorded in H₂O (A, black) and D₂O (B, red) media [20 mM HEPES and 100 mM KCl (pH or pD 7.5)] at 15 °C. The APr₆₄₈ and APg₅₄₃ forms were generated by continuous illumination with 520 and 645 nm LEDs, respectively.

APg₅₄₃-Minus-APr₆₄₈ Difference FTIR Spectra. We measured the FTIR difference spectra of AnPixJ_{GAF2} induced by illumination with an either green (520 nm) or red (645 nm) LED. The APg₅₄₃-minus-APr₆₄₈ difference spectra, which were calculated from the 250 illumination cycles in H₂O and D₂O media, are shown in Figure 6. The difference spectrum in H₂O medium showed major bands at 1703(+), 1690(−), 1676(+), 1631(−), 1602(−), 1235(+), 1220(−), and 1112(−) cm^{−1} (Figure 6A). The IR features were drastically altered by the H–D exchange (Figure 6B). All the IR bands listed above were modified in their amplitudes and/or frequencies as well as the minor bands at 1547(+), 1510(+), 1434(−), 1388(+), 1372(−), 1288(−), 1157(+), and 1146(+) cm^{−1}. The obtained APg₅₄₃-minus-APr₆₄₈ difference spectra in H₂O and D₂O are significantly different from those reported for Pfr/Pr-type phytochromes,³¹ a Pnr/Pr-type bacteriophytochrome, RbBphP3,⁷ and a Pg/Pb-type cyanobacteriochrome, TePixJ.²⁵

DISCUSSION

Features of Cyanobacteriochrome AnPixJ_{GAF2} and Phytochrome. A cyanobacteriochrome AnPixJ_{GAF2} undergoes unique green/red photoconversion.²¹ The feature is apparently different from that of red/far-red conversion in phytochromes.^{1,2} Other green sensory cyanobacteriochromes are a blue/green type,

Scheme 1. Photoconversion between APg₅₄₃ and APr₆₄₈


cyanobacteriochrome TePix], of *T. elongatus*^{18,19,25} and a green/red type, CcaS of *Synechocystis* and *Nostoc*, that regulate the chromatic adaptation.^{23,28} Therefore, the ability of cyanobacteriochromes to sense green light seems to be important for the light adaptation of cyanobacteria, which have multiple photosynthetic pigments, such as phycobilins, carotenoids, and chlorophyll *a*, to harvest light from the whole visible region.

The absorption spectrum of APg₅₄₃ in Figure 1A resembles that of the green-absorbing forms of TePix],^{18,25} although the photoconversion intermediates of the latter have not yet been identified. On the other hand, the absorption spectrum of APr₆₄₈ is quite similar to that of the Pr forms of phytochromes and bacteriophytochromes,² showing a high peak amplitude and narrow bandwidth. It is noteworthy that the photoconversions start from APr₆₄₈ and Pr to the blue and red sides, respectively. We, therefore, compare the photoconversion processes of APr₆₄₈ and Pr in detail in the following sections.

The results of this study confirm those in our previous report²¹ and indicate that each photoconversion process of AnPixJ_{GAF2} proceeds with two metastable intermediate states that can be detected 50 ns to 1 s after the laser or flash excitation. The photocycle with multiple intermediates shown in this study clearly resembles that of phytochrome.^{2,29,30} However, the features of the intermediates differ from each other. Narikawa et al. showed that AnPixJ_{GAF2} binds PCB based on an acidic urea denaturation experiment.²¹ PCB is also contained in the cyanobacterial phytochrome Cph1 of *Synechocystis*, which shows the photochromism between Pr and Pfr forms as seen in plant phytochromes.⁴³ The same PCB chromophore, thus, shows different color changes in different GAF domains. This is probably due to the interactions of identical PCBs with different protein environments inside these GAF domains.²¹ On the other hand, the shape and peak wavelength of the absorption spectrum of APr₆₄₈ are very similar to those of the Pr form of the phytochrome. This suggests that the PCB configurations of APr₆₄₈ are similar to the structure of the Pr form revealed by X-ray crystallography.⁴⁴ We summarize the green/red photocycle of APr₆₄₈ below and discuss its reaction mechanism further on the basis of the kinetic results and FTIR measurements.

Photocycle of AnPixJ_{GAF2} with Four Intermediate States. Each conversion from APg₅₄₃ or APr₆₄₈ occurred with two semistable intermediate states, as summarized in Scheme 1. We named the intermediate states in Scheme 1 according to the order of appearance and their estimated peak wavelengths. We identified intermediates G1₅₇₀ and G2₆₃₀ in the conversion from APg₅₄₃ and R1_{650–80} (which was designated as APi₆₈₀ previously in ref 21) and R2₆₁₀ in the conversion from APr₆₄₈.

The peak positions of all the intermediates, except for R1_{650–80} in Scheme 1, were different from those of the intermediates reported in the photoconversion of phytochrome, which gave peaks at 650–720 nm.^{2,29,30} APr₆₄₈ has an absorption spectrum with the highest peak amplitude and the narrowest

bandwidth in all four intermediates and APg₅₄₃. The spectral shape of APr₆₄₈ resembles that of the Pr form of the phytochrome.

Fluorescence emitted from APr₆₄₈^{*} showed a peak at 670 nm with a Stokes shift of 22 nm and a lifetime of 200 ps. The lifetime is longer than the 30–60 ps lifetimes of the Pr forms of oat and pea phytochromes, which depended on the buffer and viscosity conditions.⁴⁰ From APr₆₄₈^{*}, R1_{650–80} was formed within 100 ns, and R1_{650–80} was then converted to the second intermediate, R2₆₁₀, with a time constant of ~1 μs. R1_{650–80} may be formed either directly from APr₆₄₈^{*} with a time constant comparable to 200 ps or indirectly via some other early unidentified intermediate. Simulations in this study suggested the spectrum of R1_{650–80} (Figure 5C) has a peak at ~650 nm and a broad bandwidth extending to 680 nm. This spectrum can realize the difference spectrum, although we previously assumed R1_{650–80} to have a peak at 680 nm.²¹ We still cannot totally ignore the possibility that the difference spectrum at 100 ns is a mixture of R1_{650–80} with another earlier intermediate that has a peak at ~680 nm. Therefore, we tentatively express the intermediate as R1_{650–80} in Scheme 1. Further measurements with a shorter delay time or at low temperatures may give a clearer spectral shape of R1.

The absorption change associated with the APr₆₄₈^{*} → R1_{650–80} process can be compared to that with the Pr^{*} → lumi-R process of phytochromes or Cph1. The lumi-R intermediate form of the phytochrome has an absorption peak at 690–700 nm, at a wavelength longer than that of Pr.^{2,29,45} In Cph1, its lumi-R spectrum was assumed to have almost the same peak position as that of Pr with a lower absorption amplitude.⁴⁶ If lumi-R and R1_{650–80} share similar optical properties, then we expect similar configuration changes of the PCB pyrrole rings between lumi-R of Cph1 and R1_{650–80} of AnPixJ_{GAF2} because the spectrum of APr₆₄₈ is very similar to that of Pr of the cyanobacterial phytochrome Cph1. Then, the reaction mechanism in the first photoconversion process in AnPixJ_{GAF2} might be the *Z* ↔ *E* isomerization at the C15=C16 bond during the formation of R1_{650–80} in analogy to the interpretation of lumi-R formation.^{1,2,32}

R2₆₁₀ is assumed to have a peak at 610 nm. This indicates a blue shift from R1_{650–80} to R2₆₁₀ in the second step of the conversion. This blue shift gives a marked contrast to the red shifts between the intermediates in the Pr-to-Pfr photoconversion process of phytochromes and bacteriophytochromes.^{2,29} The blue shift of the absorption band of R2₆₁₀ suggests the localization (and/or appearance of nodal character) of the π -orbital on PCB to the smaller region due to the movement of the pyrrole ring(s) to the nonplanar orientation(s) within 1 μs.

The third step was the larger blue shift of the peak from 610 to 543 nm in the conversion from R2₆₁₀ to APg₅₄₃. This suggests the further localization of the π -orbital on PCB, probably by the further movements forming the nonplanar conformation of pyrrole rings. In AnPixJ_{GAF2}, further rotation of the A-ring may twist the plane of the pyrrole ring(s) to be out of the major molecular π -orbital plane in this step.

APg₅₄₃^{*} gave a fluorescence peak at 610 nm, which is distant from the absorption peak at 543 nm with a large Stokes shift of 70 nm, suggesting its complex excited state. APg₅₄₃^{*} fluorescence gave a lifetime of 42 ps, as shown in Table 1. This fast lifetime is comparable to the 30–60 ps lifetime of the Pr^{*} of the phytochrome.^{40,41} APg₅₄₃^{*} produced G1₅₇₀ within 50 ns. If we assume the quantum yield of the photoconversion of AnPixJ_{GAF2} to be similar to those reported for the Pr–Pfr conversion of the

phytochrome at ~ 0.1 ,^{32,43,47} then the initial photoconversion reaction, which competes with the fluorescence-emitting process, will have a reaction time constant of a few hundred picoseconds. We can, thus, assume that G1₅₇₀ is formed either directly with a time constant of a few hundred picoseconds or via another earlier intermediate that is formed with this short time constant and then converted to G1₅₇₀ within 50 ns. G1₅₇₀ turned into G2₆₃₀ with a time constant of 190 μ s, and then G2₆₃₀ turned into APr₆₄₈ with a longer time constant of 1010 μ s. The latter process was the slowest one in the whole photoconversion cycle of AnPixJ_{GAF2} and was accompanied by an increase in the absorption peak amplitude.

APg₅₄₃, G1₅₇₀, G2₆₃₀, and R2₆₁₀ states (i.e., all the states except R1_{650–80} and APr₆₄₈) are very different from the intermediates detected in the photoconversion of canonical phytochromes, which have no such blue-shifted intermediates.^{2,29,30} We can, however, find the short wavelength peaks in *Agrobacterium* Agp1 and Agp2 reconstituted with synthesized pigments.^{48,49} The absorption spectra of Agp1 and Agp2 with “15Zs” and “15Es” locked BV, in which the C15=C16 bond was fixed in the Z/E configuration and the C14–C15 bond in the *syn* conformation, showed peaks at ~ 430 nm. No intermediate of AnPixJ_{GAF2} showed such a blue peak. On the other hand, Agp2 reconstituted with a “5Za” synthetically locked chromophore, which is a BV derivative with the C4=C5 bond fixed in the Z configuration and the C5–C6 bond in an *anti* conformation, showed the absorption peak at 640 nm. This peak wavelength is close to that of G2₆₃₀ of AnPixJ_{GAF2}. The configuration of PCB in G2₆₃₀ might resemble this conformation, although further studies with AnPixJ should be conducted to confirm it.

Structural Changes of the Pigment and Protein in the Photoconversion of AnPixJ_{GAF2}. In the reaction of Cph1 and phytochromes, several reports proposed the Z \leftrightarrow E isomerization at the C15=C16 bond (of the D-ring) in the conversion from Pr to Pfr.^{1,2,32} X-ray structural analysis of α -phycoerythrocyanin also suggested the Z \leftrightarrow E isomerization at the C15=C16 bond of phycoviolobilin to be responsible for the shift of the peak from 500 nm (15E) to 580 nm (15Z).⁵⁰ A recent NMR study of the cyanobacterial phytochrome of *Synechococcus* OSB' proposed a different mechanism, namely, that the isomerization of the A-ring of PCB is responsible for the conversion from the Pr form to the Pfr form.⁵¹ Although no comparable experimental results exist for AnPixJ at present, we can assume the isomerization (rotation) of the pyrrole ring(s) on either side edge of PCB to be responsible for the conversion to the shorter wavelength. APr₆₄₈ showed the sequential blue shifts in the conversion to APg₅₄₃, while the Pr of the phytochrome or bacteriophytochrome with a similar absorption spectrum shows the red shift in the photoconversion to Pfr. Therefore, the changes in configuration of the PCB chromophores in AnPixJ_{GAF2} and phytochromes seem to differ in the chromophore–protein interactions.

The APg₅₄₃-minus-APr₆₄₈ difference FTIR spectrum showed features (Figure 6) that were significantly different from those reported for the photoconversion of phytochromes, TePixJ, Agp1, and RbBphP3.^{7,25,31} Although the information obtained in this study is limited, we tentatively assigned the IR peaks/troughs as described below by comparing the spectrum with that of the Pr/Pfr-type cyanobacterial phytochrome, Cph1, which also contains PCB.³¹ It is known that the Pfr/Pr-type phytochromes present a marker band of the Pfr state at 1650–1656 cm^{-1} . In the Pfr-minus-Pr spectrum of Cph1, the most prominent band was detected at 1653 cm^{-1} . This band was less sensitive to the H–D

exchange as well as to the specific isotope labeling of the PCB itself and thus can be assigned primarily to amide I of the α -helical structure of the protein.³¹ In the amide I region of AnPixJ_{GAF2} spectra, no such marker band was evident. We, therefore, assumed that the IR feature at 1666(+)/1631(–) cm^{-1} , which was less sensitive to H–D exchange, reflects the changes in the amide I and/or chromophore C=C vibrational modes. A remarkable negative band in the amide I region was also resolved in the FTIR spectrum of the domain-truncated Agp1.⁵² The Pfr-minus-Pr difference spectrum of the Agp1-M15 protein, which is composed of the chromophore module (PLD, GAF, and PHY domains) and lacks the His-kinase module, shows a negative band at 1635 cm^{-1} . The apparent similarity of the negative band in the amide I region (1631 cm^{-1} in AnPixJ_{GAF2} and 1635 cm^{-1} in Agp1-M15) might represent a similarity in protein architecture between APr₆₄₈ of AnPixJ_{GAF2} and Pr of Agp1-M15.

In the carbonyl C=O stretch region of the Pfr-minus-Pr difference spectrum of Cph1, the bands at 1736(–)/1724(+) and 1704(–) cm^{-1} were assigned to the carbonyl groups of PCB pyrrole ring-A and ring-D, respectively, on the basis of the effects of H–D exchange and specific isotope labeling on PCB. In AnPixJ_{GAF2}, IR bands at 1730(–), 1703(+), 1690(–), and 1676(–) cm^{-1} , which were sensitive to H–D exchange, could have arisen from the vibrational modes of the protonated and/or hydrogen-bonded chemical groups. On the other hand, in an analogy to the cases in Cph1³¹ and/or phyA,⁵² the 1730(–), 1703(+), and 1602(–) cm^{-1} bands could be assigned to the structural changes in PCB ring-A and ring-D and the B–C bridge, respectively. The 1690(–)/1676(+) cm^{-1} feature might be assigned to the propionate of PCB. In a lower-frequency region, the prominent IR features at 1235(+)/1220(–) and 1112(–) cm^{-1} , which were apparently lost in the D₂O medium, seem to be assigned to the C–N vibrational modes of PCB according to ref 31 in which the IR bands in this region were tentatively assigned to the C–N mode of PCB in Cph1. In the case of Agp1-M15, a band shift at 1244(–)/1233(+) cm^{-1} has been resolved in the Pfr-minus-Pr spectrum.⁵³ These trough/peak positions are close to those of AnPixJ_{GAF2} [1235(+)/1220(–) cm^{-1}], but the direction of the shift is opposite. These results indicate that the chromophore–protein interaction in AnPixJ_{GAF2} is largely different from that in phytochromes, in agreement with the prediction from the analysis of the visible spectral changes. It was also reported that B-deprotonated tetrapyrrole shows a downshift in the IR band at 1224 cm^{-1} (A-ring N–H in-plane bending) by DFT calculations.⁵³ Thus, the band pair at 1235(+)/1220(–) cm^{-1} may be derived from the change in protonation state between APr₆₄₈ and APg₅₄₃, although it is difficult to assign at present. It is noteworthy that deprotonation and protonation take place, as proposed for the Pr \rightarrow Pfr conversion of phytochromes,³⁵ during the photoconversion of AnPixJ_{GAF2}.

We tested the structural difference between phytochromes and AnPixJ_{GAF2} on the basis of their amino acid sequences, shown in Figure 1D on the structure of the former. The residues shown in Figure 1D are shown in Figure 1E in the ball-and-stick model to elucidate their location on the known structure of the Pr form of SyCph1. The residues, which are homologous to those of AnPixJ_{GAF2}, are colored red and yellow, and those that are not conserved in AnPixJ_{GAF2} and are in contact with PCB are colored blue. PCB drawn in the stick model is colored white. The image shows that the residues in contact with PCB (within 5 Å) are

rather unconserved (colored blue) except for a few conserved residues that are colored yellow. The majority of conserved residues, which are colored red, are far from PCB. The distributions of conserved and unconserved residues, therefore, seem to support the conclusions from the FTIR studies and intermediate analyses in this study that assessed the interaction between the PCB chromophore and protein to be significantly different from that in phytochromes. The unique unconserved residues in contact with PCB seem to represent the unique photoconversion mechanism of AnPixJ_{GAF2}. Nature, thus, has created different color sensory systems in cyanobacteriochromes and phytochromes by almost fully modifying the amino acid residues just in contact with the pigment within the homologous outer shell of the GAF domains.

Photoconversion Mechanism of AnPixJ_{GAF2}. AnPixJ_{GAF2} undergoes the photoconversion cycle between its red and green forms with four intermediate states as shown in Scheme 1, i.e., two intermediate states in each direction. The feature with the stepwise peak shifts in each direction resembles that of phytochromes and bacteriophytochromes and the other photoreceptors in which isomerization of the pigment takes place.¹² The phytochrome photoconversion has been assumed to accompany the *Z* ↔ *E* isomerization at a C15=C16 bond. The movement of similar pyrrole ring(s) (presumably the D-ring) may also occur in APr₆₄₈. The peak wavelengths of R2₆₁₀, G1₅₇₀, and G2₆₃₀ in the AnPixJ_{GAF2} cycle, however, are apparently different from those of intermediates in the photocycles of phytochromes. To realize the shorter wavelength peaks of intermediates, different molecular mechanisms, such as unique movements and/or rotations of pyrrole A- or D-rings, may occur in AnPixJ_{GAF2}. The reconstitution of pigments with locked configurations, like that in Agp1 and Agp2,^{48,49} will be useful in understanding its photoconversion mechanism.

The FTIR difference spectrum of AnPixJ_{GAF2}, which is significantly different from those of phytochromes, suggests the interaction between the PCB chromophore and AnPixJ_{GAF2} proteins to be somewhat different from that of phytochromes. The multiple vibrational modes were sensitive to the photoconversion and to H–D exchange, suggesting the high accessibility of protein and chromophore side bands to water in AnPixJ_{GAF2}, although the PCB configurations in the APr₆₄₈ and Pr forms appear to be similar according to their similar absorption spectra. This may suggest a slightly exposed location of PCB in AnPixJ_{GAF2}. The homology of the amino acid residues visualized in Figure 1E clearly indicates that the outer shell structure of the GAF domain is well-conserved, while the chromophore-surrounding residues, including the ligand residues, are significantly modified. The unique feature of the photocycle of AnPixJ_{GAF2}, thus, seems to be realized by the variations of the residues that surround and ligate the PCB pigment. The high degree of sensitivity to H–D exchange also suggests the shift of the PCB location closer to the outer aqueous phase from that in Figure 1E.

AnPixJ_{GAF2} was recently crystallized,⁵⁴ and its structure in the APr₆₄₈ form will be revealed soon. A preliminary analysis indicated that PCB shows a configuration similar to that of Cph1 in the Pr form in a shifted binding position (R. Narikawa, N. Muraki, T. Shiba, G. Kurisu, and M. Ikeuchi, unpublished result), which agrees with the conclusion of this study. Analyses based on the structure will further advance our understanding of the photoconversion mechanism of green/red or blue/green sensing by cyanobacteriochromes and may also extend our

understanding of the photoconversion mechanism of phytochromes. We will be able to design the proteins with a variety of colors and reaction modes with this knowledge.

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Funding Sources

This work has been supported by grant-in-aid for challenging exploratory research to S.I. (23655196) from MEXT.

ACKNOWLEDGMENT

We thank Drs. Peter R. Rich and Santiago Garcia (University College London) for their kind assistance with the FTIR measurement and Dr. Masahiro Ishiura in the Center for Gene Research of Nagoya University and Dr. Kosuke Maki in the Department of Physics of Nagoya University for their kind support of the research.

ABBREVIATIONS

APg₅₄₃ and APr₆₄₈, green- and red-absorbing forms of the second GAF domain of AnPixJ, respectively; BV, biliverdin; EADS, evolution-associated difference spectrum; FTIR, Fourier transform infrared spectroscopy; GAF, cGMP-phosphodiesterase/*Anabaena* adenyllyl-cyclase/*E. coli* formate hydrogen lyase transcription activator; PCB, phycocyanobilin; Pr and Pfr, red- and far-red-absorbing forms of the phytochrome, respectively.

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